

**ASSEMBLY BILL**

**No. 1370**

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**Introduced by Assembly Member Matthews**

February 22, 2005

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An act to amend Section 1209 of the Business and Professions Code, relating to clinical laboratories.

LEGISLATIVE COUNSEL'S DIGEST

AB 1370, as introduced, Matthews. Clinical laboratory director: pharmacists.

Existing law provides for the licensure and regulation of clinical laboratories and their personnel by the State Department of Health Services and makes a violation of those provisions a crime. Existing law defines various terms for this purpose, including "laboratory director."

This bill would include a pharmacist within the definition of laboratory director if the clinical laboratory test or examination is a routine patient assessment procedure, as defined. Because a failure of such a pharmacist to comply with this particular limitation or with the provisions regulating clinical laboratories and their personnel would be punishable as a crime, the bill would impose a state-mandated local program.

The California Constitution requires the state to reimburse local agencies and school districts for certain costs mandated by the state. Statutory provisions establish procedures for making that reimbursement.

This bill would provide that no reimbursement is required by this act for a specified reason.

Vote: majority. Appropriation: no. Fiscal committee: yes.  
State-mandated local program: yes.

*The people of the State of California do enact as follows:*

SECTION 1. Section 1209 of the Business and Professions Code is amended to read:

1209. (a) As used in this chapter, “laboratory director” means any person who is a duly licensed physician and surgeon, or is licensed to direct a clinical laboratory under this chapter and who substantially meets the laboratory director qualifications under CLIA for the type and complexity of tests being offered by the laboratory. *“Laboratory director” also means a pharmacist if the clinical laboratory test or examination is a routine patient assessment procedure, as defined in Section 4052.1.* The laboratory director, if qualified under CLIA, may perform the duties of the technical consultant, technical supervisor, clinical consultant, general supervisor, and testing personnel, or delegate these responsibilities to persons qualified under CLIA. If the laboratory director reappoints performance of those responsibilities or duties, he or she shall remain responsible for ensuring that all those duties and responsibilities are properly performed.

(b) (1) The laboratory director is responsible for the overall operation and administration of the clinical laboratory, including administering the technical and scientific operation of a clinical laboratory, the selection and supervision of procedures, the reporting of results, and active participation in its operations to the extent necessary to assure compliance with this act and CLIA. He or she shall be responsible for the proper performance of all laboratory work of all subordinates and shall employ a sufficient number of laboratory personnel with the appropriate education and either experience or training to provide appropriate consultation, properly supervise and accurately perform tests, and report test results in accordance with the personnel qualifications, duties, and responsibilities described in CLIA and this chapter.

(2) Where a point-of-care laboratory testing device is utilized and provides results for more than one analyte, the testing personnel may perform and report the results of all tests ordered for each analyte for which he or she has been found by the laboratory director to be competent to perform and report.

(c) As part of the overall operation and administration, the laboratory director of a registered laboratory shall document the

adequacy of the qualifications (educational background, training, and experience) of the personnel directing and supervising the laboratory and performing the laboratory test procedures and examinations. In determining the adequacy of qualifications, the laboratory director shall comply with any regulations adopted by the department that specify the minimum qualifications for personnel, in addition to any CLIA requirements relative to the education or training of personnel.

(d) As part of the overall operation and administration, the laboratory director of a licensed laboratory shall do all of the following:

(1) Ensure that all personnel, prior to testing biological specimens, have the appropriate education and experience, receive the appropriate training for the type and complexity of the services offered, and have demonstrated that they can perform all testing operations reliably to provide and report accurate results. In determining the adequacy of qualifications, the laboratory director shall comply with any regulations adopted by the department that specify the minimum qualifications for, and the type of procedures that may be performed by, personnel in addition to any CLIA requirements relative to the education or training of personnel. Any regulations adopted pursuant to this section that specify the type of procedure that may be performed by testing personnel shall be based on the skills, knowledge, and tasks required to perform the type of procedure in question.

(2) Ensure that policies and procedures are established for monitoring individuals who conduct preanalytical, analytical, and postanalytical phases of testing to assure that they are competent and maintain their competency to process biological specimens, perform test procedures, and report test results promptly and proficiently, and, whenever necessary, identify needs for remedial training or continuing education to improve skills.

(3) Specify in writing the responsibilities and duties of each individual engaged in the performance of the preanalytic, analytic, and postanalytic phases of clinical laboratory tests or examinations, including which clinical laboratory tests or examinations the individual is authorized to perform, whether supervision is required for the individual to perform specimen processing, test performance, or results reporting, and whether

1 consultant, supervisor, or director review is required prior to the  
2 individual reporting patient test results.

3 (e) The competency and performance of staff of a licensed  
4 laboratory shall be evaluated and documented by the laboratory  
5 director, or by a person who qualifies as a technical consultant or  
6 a technical supervisor under CLIA depending on the type and  
7 complexity of tests being offered by the laboratory.

8 (1) The procedures for evaluating the competency of the staff  
9 shall include, but are not limited to, all of the following:

10 (A) Direct observations of routine patient test performance,  
11 including patient preparation, if applicable, and specimen  
12 handling, processing, and testing.

13 (B) Monitoring the recording and reporting of test results.

14 (C) Review of intermediate test results or worksheets, quality  
15 control records, proficiency testing results, and preventive  
16 maintenance records.

17 (D) Direct observation of performance of instrument  
18 maintenance and function checks.

19 (E) Assessment of test performance through testing previously  
20 analyzed specimens, internal blind testing samples, or external  
21 proficiency testing samples.

22 (F) Assessment of problem solving skills.

23 (2) Evaluation and documentation of staff competency and  
24 performance shall occur at least semiannually during the first  
25 year an individual tests biological specimens. Thereafter,  
26 evaluations shall be performed at least annually unless test  
27 methodology or instrumentation changes, in which case, prior to  
28 reporting patient test results, the individual's performance shall  
29 be reevaluated to include the use of the new test methodology or  
30 instrumentation.

31 (f) The laboratory director of each clinical laboratory of an  
32 acute care hospital shall be a physician and surgeon who is a  
33 qualified pathologist, except as follows:

34 (1) If a qualified pathologist is not available, a physician and  
35 surgeon or a clinical laboratory bioanalyst qualified as a  
36 laboratory director under subdivision (a) may direct the  
37 laboratory. However, a qualified pathologist shall be available  
38 for consultation at suitable intervals to ensure high quality  
39 service.

1 (2) If there are two or more clinical laboratories of an acute  
2 care hospital, those additional clinical laboratories that are  
3 limited to the performance of blood gas analysis, blood  
4 electrolyte analysis, or both may be directed by a physician and  
5 surgeon qualified as a laboratory director under subdivision (a),  
6 irrespective of whether a pathologist is available.

7 As used in this subdivision, a qualified pathologist is a  
8 physician and surgeon certified or eligible for certification in  
9 clinical or anatomical pathology by the American Board of  
10 Pathology or the American Osteopathic Board of Pathology.

11 (g) Subdivision (f) does not apply to any director of a clinical  
12 laboratory of an acute care hospital acting in that capacity on or  
13 before January 1, 1988.

14 SEC. 2. No reimbursement is required by this act pursuant to  
15 Section 6 of Article XIII B of the California Constitution because  
16 the only costs that may be incurred by a local agency or school  
17 district will be incurred because this act creates a new crime or  
18 infraction, eliminates a crime or infraction, or changes the  
19 penalty for a crime or infraction, within the meaning of Section  
20 17556 of the Government Code, or changes the definition of a  
21 crime within the meaning of Section 6 of Article XIII B of the  
22 California Constitution.